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# Transcriptome-wide association study of breast cancer risk by estrogen-receptor status

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 HEBON Investigators<sup>120</sup> | BCFR Investigators<sup>121</sup> | OCGN Investigators<sup>122</sup> |  
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 Sara Lindström<sup>139,140</sup> | Wing-Yee Lo<sup>44,45</sup> | Jennifer T. Loud<sup>107</sup> | Jan Lubiński<sup>29</sup> |  
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 Maria E. Martinez<sup>145,146</sup> | Laura Matricardi<sup>147</sup> | Tabea Maurer<sup>63</sup> |  
 Dimitrios Mavroudis<sup>148</sup> | Lesley McGuffog<sup>13</sup> | Alfons Meindl<sup>149</sup> | Usha Menon<sup>150</sup> |  
 Kyriaki Michailidou<sup>13,151</sup> | Pooja M. Kapoor<sup>25,152</sup> | Austin Miller<sup>153</sup> |  
 Marco Montagna<sup>147</sup> | Fernando Moreno<sup>54</sup> | Lidia Moserle<sup>147</sup> |  
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 Nisha Pradhan<sup>129</sup> | Karolina Prajzendanc<sup>29</sup> | Nadege Presneau<sup>84</sup> | Kevin Punie<sup>157</sup> |  
 Katri Pylkäs<sup>168,169</sup> | Paolo Radice<sup>170</sup> | Johanna Rantala<sup>171</sup> |  
 Muhammad Usman Rashid<sup>111,172</sup> | Gad Rennert<sup>8</sup> | Harvey A. Risch<sup>173</sup> |  
 Mark Robson<sup>160</sup> | Atocha Romero<sup>174</sup> | Emmanouil Saloustros<sup>175</sup> |  
 Dale P. Sandler<sup>176</sup> | Catarina Santos<sup>165</sup> | Elinor J. Sawyer<sup>177</sup> |  
 Marjanka K. Schmidt<sup>50,178</sup> | Daniel F. Schmidt<sup>101,179</sup> | Rita K. Schmutzler<sup>71,180</sup> |  
 Minouk J. Schoemaker<sup>99</sup> | Rodney J. Scott<sup>181,182,183</sup> | Priyanka Sharma<sup>184</sup> |  
 Xiao-Ou Shu<sup>7</sup> | Jacques Simard<sup>185</sup> | Christian F. Singer<sup>19</sup> | Anne-Bine Skytte<sup>186</sup> |  
 Penny Soucy<sup>185</sup> | Melissa C. Southey<sup>187,188</sup> | John J. Spinelli<sup>189,190</sup> |  
 Amanda B. Spurdle<sup>163</sup> | Jennifer Stone<sup>101,191</sup> | Anthony J. Swerdlow<sup>99,192</sup> |

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**Nadine Tung**<sup>204</sup> | **Celine M. Vachon**<sup>205</sup> | **Christi J. van Asperen**<sup>206</sup> |  
**Ans M. W. van den Ouweland**<sup>207</sup> | **Elizabeth J. van Rensburg**<sup>208</sup> | **Ana Vega**<sup>26,30,31</sup> |  
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**Camilla Wendt**<sup>144</sup> | **Robert Winqvist**<sup>168,169</sup> | **Xiaohong R. Yang**<sup>64</sup> |  
**Drakoulis Yannoukakos**<sup>93</sup> | **Argyrios Ziogas**<sup>12</sup> | **Roger L. Milne**<sup>100,101,187</sup> |  
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### Abstract

Previous transcriptome-wide association studies (TWAS) have identified breast cancer risk genes by integrating data from expression quantitative loci and genome-wide association studies (GWAS), but analyses of breast cancer subtype-specific associations have been limited. In this study, we conducted a TWAS using gene expression data from GTEx and summary statistics from the hitherto largest GWAS meta-analysis conducted for breast cancer overall, and by estrogen receptor subtypes (ER+ and ER–). We further compared associations with ER+ and ER– subtypes, using a case-only TWAS approach. We also conducted multigene conditional analyses in regions with multiple TWAS associations. Two genes, *STXBP4* and *HIST2H2BA*, were specifically associated with ER+ but not with ER– breast cancer. We further identified 30 TWAS-significant genes associated with overall breast cancer risk, including four that were not identified in previous studies. Conditional analyses identified single independent breast-cancer gene in three of six regions harboring multiple TWAS-significant genes. Our study provides new information on breast cancer genetics and biology, particularly about genomic differences between ER+ and ER– breast cancer.

### KEYWORDS

breast cancer subtype, causal gene, GWAS, TWAS

## 1 | INTRODUCTION

Breast cancer is the most common malignancy among women worldwide (Bray et al., 2018). The disease has a strong inherited component (Beggs & Hodgson, 2009); linkage studies have identified infrequent mutations in *BRCA1/2* (Easton et al., 2007; Seal et al., 2006; Turnbull et al., 2010) and genome-wide association studies (GWAS) have identified 177 susceptibility loci to date (Michailidou et al., 2017). However, these GWAS-discovered variants explain only 18% of the familial relative risk of breast cancer. Moreover, the causal mechanism driving GWAS associations remains largely unknown, as many variants are located in noncoding or intergenic regions, and are not in strong linkage disequilibrium (LD) with known protein-coding variants (Beggs & Hodgson, 2009; Michailidou et al., 2015).

Breast cancer is a heterogeneous disease consisting of several well-established subtypes. One of the most important markers of breast cancer subtypes is estrogen receptor (ER) status. ER+ and ER– tumors differ in etiology (X. R. Yang, Chang-Claude, et al., 2011), genetic

predisposition (Mavaddat, Antoniou, Easton, & Garcia-Closas, 2010), and clinical behavior (Blows et al., 2010). ER– tumor occurs more often among younger women, and patients are more likely to carry *BRCA1* pathogenic variants (Atchley et al., 2008; Garcia-Closas et al., 2013). ER– tumor also has worse short-term prognosis. Among the 177 GWAS-identified breast cancer-associated single nucleotide polymorphisms (SNPs), around 50 are more strongly associated with ER+ disease and 20 are more strongly associated with ER– disease (Michailidou et al., 2017; Milne et al., 2017).

SNPs associated with complex traits are more likely to be in regulatory regions than in protein-coding regions, and many of these SNPs are also associated with expression levels of nearby genes (Nicolae et al., 2010). For example, breast cancer GWAS-identified variants at 6q25.1 regulate *ESR1*, but also coregulate other local genes such as *RMND1*, *ARMT1*, and *CCDC170* (Dunning et al., 2016, p. 1). These results suggest that by integrating genotype, phenotype, and gene expression, we can identify novel trait-associated genes and understand biological mechanisms. However, due to costs and tissue

availability, acquiring GWAS and gene expression data for the same set of individuals remains challenging.

A recently published approach, referred to as transcriptome-wide association study (TWAS; Gamazon et al., 2015; Gusev et al., 2016), overcomes these difficulties by using a relatively small set of reference individuals for whom both gene expression and SNPs have been measured to impute the *cis*-genetic component of expression for a much larger set of individuals from their GWAS summary statistics. The association between the predicted gene expression and traits can then be tested. This method has been shown to have greater power relative to GWAS; and has identified 1,196 trait-associated genes across 30 complex traits in a recently performed multitissue TWAS (Mancuso et al., 2017).

To date, three TWAS of breast cancer have been conducted (Gao, Pierce, Olopade, Im, & Huo, 2017; Hoffman et al., 2017; Wu et al., 2018). A fourth study linked expression quantitative loci (eQTL) data across multiple tissues and breast cancer GWAS results using EUGENE, a statistical approach that sums evidence for association with disease across eQTLs regardless of directionality. That study then tested EUGENE-significant genes using a TWAS statistic, which does take directionality into account (Ferreira et al., 2019). The two earliest TWAS used GWAS data from the National Cancer Institute's "Up for a Challenge" competition, which included data from 12,100 breast cancer cases (of which 3,900 had ER– disease) and 11,400 controls, as well as eQTL data from breast tissue and whole blood from the GTEx and DGN projects (Gao et al., 2017; Hoffman et al., 2017). The subsequent TWAS by Wu et al. (2018) and the EUGENE analysis by Ferreira et al. (2019) used results from a much larger GWAS conducted by the Breast Cancer Association Consortium (BCAC), which included 122,977 cases (of which 21,468 had ER– disease) and 105,974 controls. Together, these four studies have identified 59 genes whose predicted expression levels are associated with risk of overall breast cancer, and five associated with risk of ER– disease. Of these 64 genes, 30 are at loci not previously identified by breast cancer GWAS.

These previous TWAS largely focused on overall breast cancer risk. Analyses of ER– disease either were conducted using a small sample size (Gao et al., 2017) or did not scan all genes using a directional TWAS approach (Ferreira et al., 2019). Moreover, none of the previous analyses considered ER+ disease specifically or examined differences in association between predicted gene expression and ER+ versus ER– disease.

The interpretation of TWAS results is not straightforward (Wainberg et al., 2019). Specifically, TWAS statistic by itself cannot distinguish between a mediated effect (SNPs influence breast cancer risk by changing the

expression of the tested gene), pleiotropy (SNPs associated with gene expression also influence breast cancer risk through another mechanism), or colocalization (SNPs associated with gene expression are in LD with other SNPs that influence breast cancer risk through another mechanism). Previous studies have conducted limited sensitivity analyses (e.g., Wu et al., 2018 and Ferreira et al., 2019 conditioned the TWAS tests on lead GWAS SNPs), but the genetic architecture at TWAS-identified loci remains largely unclear.

In the current analysis, we complement previous work by conducting a TWAS for overall breast cancer and for ER+ and ER– subtypes. We also applied a case-only TWAS test to identify predicted transcript levels that were differentially associated with ER+ and ER– disease. We conducted expanded sensitivity analyses, conditioning on multiple TWAS-significant genes in a region to account for possible confounding due to LD (colocalization). We chose to focus on the expression of normal breast tissue of European ancestry women to maximize specificity and identify good targets for near-term follow-up experiments in mammary cells. One advantage of using a biologically relevant tissue is that it both increases the *a priori* plausibility of observed associations and increases the likelihood that genes with observed associations will be expressed and influence tumor development in cells from the target tissue. We have reproduced previous results (Ferreira et al., 2019; Wu et al., 2018) and provided evidence regarding the independent associations of multiple genes in regions containing one or more TWAS-significant genes. We also identified genes with subtype-specific associations, highlighting different biological mechanisms likely underlying the disease subtypes.

## 2 | MATERIAL AND METHODS

### 2.1 | Gene expression reference panel

The transcriptome and high-density genotyping data used to build the gene expression model (reference panel) were retrieved from GTEx (GTEx Consortium, 2015), a consortium collected high-quality gene expression RNA-seq data across 44 body sites from 449 donors, and genome-wide genetic information. For the current study, we included 67 women of European ancestry who provided normal breast mammary tissues. RNA samples extracted from tissues were sequenced to generate data on 12,696 transcripts. Genomic DNA samples were genotyped using Illumina OMNI 5 M or 2.5 M arrays, processed with a standard GTEx protocol. Briefly, SNPs with call rates <98%, with differential missingness between the two array experiments (5 M or 2.5 M), with Hardy-Weinberg equilibrium  $p < 10^{-6}$  or showing batch

effects were excluded. The genotypes were then imputed to the Haplotype Reference Consortium reference panel (McCarthy et al., 2016) using Minimac3 for imputation and SHAPEIT for pre-phasing (Delaneau, Marchini, & Zagury, 2011; Howie, Donnelly, & Marchini, 2009). Only SNPs with high imputation quality ( $r^2 \geq .8$ ), minor allele frequency (MAF)  $\geq 0.05$ , and were included in the HapMap Phase 2 version were used to build the expression prediction models.

## 2.2 | Breast cancer meta-GWAS data

The GWAS breast cancer summary-level data were mainly provided by the Breast Cancer Association Consortium (BCAC; Michailidou et al., 2017), as well as the Consortium of Investigators of Modifiers of *BRCA1/2* (CIMBA). BCAC conducted the largest breast cancer meta-GWAS to date (referred as the overall breast cancer GWAS analysis). The BCAC included 122,977 cases and 105,974 controls of European ancestry. Among these, 46,785 cases and 42,892 controls were genotyped using the Illumina iSelect genotyping array (iCOGS) on 211,155 SNPs; and 61,282 cases and 45,494 controls were genotyped using the Illumina OncoArray on 570,000 SNPs (Yovel, Franz, Stilz, & Schnitzler, 2008). The study also included data from 11 other GWAS on 14,910 cases and 17,588 controls. Genetic data for all individual participating studies were imputed to the 1000 Genomes Project Phase 3 v5 EUR reference panel. Logistic regression was fitted to estimate per-allele odds ratios (ORs), adjusting for country and top principal components (PCs). Inverse variance fixed-effect meta-analysis was used to combine the genetic association for breast cancer risk across studies (Milne et al., 2017). In CIMBA, genotypes were generated by the Illumina OncoArray and imputed to the 1000 Genomes Project Phase 3 v5 EUR reference panel (Amos et al., 2016). A retrospective cohort analysis framework was adopted to estimate per-allele hazard ratios (HRs), modelling time-to-breast-cancer and stratified by country, Ashkenazi Jewish origin and birth cohort (Antoniou et al., 2005; Barnes et al., 2012). Fixed-effect meta-analysis (Willer, Li, & Abecasis, 2010) was performed to combine results across genotyping initiatives within the two consortia, assuming that the OR and HR estimates had roughly the same underlying relative risk. We restricted subsequent analyses to SNPs with an imputation  $r^2 > .3$ , and an MAF  $> 0.005$  across all platforms were included in the analysis (approximately 11.5 M).

For the ER+ subtype, meta-GWAS summary data based on 69,501 ER+ cases and 105,974 controls (part of the overall breast cancer samples) were included and analyzed (Mavaddat et al., 2015). For the ER– subtype,

meta-GWAS summary data based on 21,468 ER– cases and 105,974 controls from the BCAC were combined with 9,414 additional *BRCA1* mutation-positive cases and 9,494 *BRCA1* mutation-positive controls from CIMBA (Milne et al., 2017).

To distinguish different genetic signals between ER+ and ER– subtypes, we further retrieved GWAS summary-level data on a case-only GWAS, which compared ER+ patients (sample size: 23,330 in iCOGs and 44,746 in OncoArray) to ER– patients (sample size: 5,479 and 11,856; Milne et al., 2017). Logistic regression was performed to test the association between genetic variants with known ER status in the two studies separately, adjusting for substudy and top PCs for iCOGs, and patients' countries and top PCs for OncoArray. Results were then combined using a fixed-effect meta-analysis.

## 2.3 | Constructing expression weights

Before constructing the expression model (using GTEx data, regress gene expression on SNPs), we set several criteria to select eligible candidate genes for inclusion in the model (from the total 12,696 transcripts). We used a REML algorithm implemented in GCTA to estimate the *cis* (500 base-pair window surrounding transcription start site) SNP-heritability ( $cis-h_g^2$ ) for each transcript expression (Cai et al., 2014; J. Yang et al., 2010). Only genes with significant heritability (nominal  $p \leq .01$ ) were included in the subsequent model construction (J. Yang, Lee, Goddard, & Visscher, 2011). The  $p$  values for null hypotheses  $cis-h_g^2 = 0$  were computed using a likelihood ratio test. To account for population stratification, 20 PCs were always included as fixed effects. Consistent with previous research (ENCODE Project Consortium, 2012; J. Yang et al., 2010), we observed strong evidence for  $cis-h_g^2$  on many genes (significantly non-zero for 1,355 genes).

We then constructed linear genetic predictors of gene expression for these genes. We performed five models: Bayesian Sparse Linear Mixed model, Best Linear Unbiased Predictor model, Elastic-net regression (with mixing parameter of 0.5), LASSO regression and Single best eQTL model. We used a fivefold cross-validation strategy to validate each model internally. Only genes with good model performance, corresponding to a prediction  $r^2$  value (the square of the correlation between predicted and observed expression) of at least 1% (0.10 correlation) in at least one of the five models, were included in subsequent TWAS analyses. The weights were chosen from the best performed model out of the five models. We adopted this additional filter to improve the interpretability and specificity of results: significant TWAS results based on models with little or no predictive ability likely result from pleiotropy or

**TABLE 1** Genes significantly associated with the overall breast cancer, estrogen receptor positive and negative subtypes, and estrogen receptor status, as identified by TWAS

Cytoband	Gene	Chromosome: Position (start–end)	Number of SNPs	Heritability	Cross validation $r^2$	TWAS $p$ values			
						Overall vs. controls	ER+ vs. controls	ER– vs. controls	ER+ vs. ER–
p11.2	HIST2H2BA	1: 120906028–120915073	77	0.10	0.04	<b>3.1E–30</b>	<b>2.0E–32</b>	3.6E–01	<b>1.4E–07</b>
q21.1	NUDT17	1: 145586804–145589439	110	0.07	0.03	<b>1.7E–09</b>	<b>8.8E–10</b>	2.4E–01	3.3E–03
q33.1	ALS2CR12	2: 202152994–202222121	363	0.35	0.03	<b>2.2E–11</b>	<b>8.4E–07</b>	<b>2.5E–06</b>	8.8E–01
	CASP8	2: 202098166–202152434	371	0.32	0.15	<b>1.8E–07</b>	<b>8.2E–06</b>	2.9E–04	7.6E–01
q25.31	LINC00886	3: 1156465135–156534851	452	0.17	0.04	<b>4.7E–05</b>	1.9E–04	3.2E–03	7.3E–01
p16.3	MAEA	4: 1283639–1333925	374	0.42	0.23	<b>3.9E–05</b>	1.6E–04	2.3E–01	2.3E–01
q14.2	ATG10	5: 81267844–81551958	520	0.44	0.26	<b>1.9E–10</b>	<b>2.4E–10</b>	2.7E–01	2.4E–02
	ATP6AP1L	5: 81575281–81682796	467	0.62	0.51	<b>2.3E–07</b>	<b>3.3E–06</b>	2.9E–02	2.3E–01
q14.1	RP11–250B2.5	6: 81176675–81178797	402	0.12	0.08	<b>6.9E–07</b>	7.3E–03	5.9E–03	8.2E–01
q22.33	RP11–73O6.3	6: 130454555–1130465515	557	0.33	0.11	<b>4.5E–12</b>	<b>1.2E–06</b>	<b>5.2E–09</b>	5.5E–02
	L3MBTL3	6: 130334844–130462594	609	0.31	0.23	<b>8.5E–14</b>	<b>3.6E–08</b>	<b>1.8E–10</b>	1.4E–01
p15.1	GDI2	10: 5807186–5884095	727	0.29	0.01	<b>4.8E–07</b>	<b>3.1E–07</b>	4.0E–01	2.1E–01
p15.5	MRPL23-AS1	11: 2004467–2011150	449	0.34	0.09	<b>7.4E–06</b>	1.4E–04	2.9E–01	2.2E–01
q13.2	RP11–554A11.9	11: 68923378–68927220	428	0.23	0.08	<b>1.5E–06</b>	4.9E–04	7.1E–03	6.9E–01
q15	NUP107	12: 69082742–69136785	538	0.27	0.05	<b>6.1E–06</b>	<b>8.2E–07</b>	4.3E–01	1.0E–01
q24.3	ULK3	15: 75128457–75135538	286	0.19	0.15	<b>3.9E–05</b>	<b>2.1E–05</b>	2.8E–02	5.2E–01
	MAN2C1	15: 75648133–75659986	226	0.37	0.34	<b>7.4E–07</b>	1.7E–04	1.8E–03	6.3E–01
	CTD–2323K18.1	15: 75819491–75893546	226	0.36	0.19	<b>7.5E–07</b>	3.9E–04	2.3E–03	3.1E–01
q21.31	HSD17B1P1	17: 40698782–40700724	245	0.09	0.07	<b>1.4E–06</b>	1.8E–03	6.4E–03	3.9E–01
q21.32	LRRC37A4P	17: 43578685–43623042	149	0.34	0.35	<b>3.1E–10</b>	<b>1.4E–06</b>	2.2E–02	8.0E–01
	CRHR1-IT1	17: 43697694–43725582	87	0.35	0.41	<b>2.9E–10</b>	<b>1.8E–06</b>	2.7E–02	8.2E–01
	CRHR1	17: 43699267–43913194	86	0.09	0.18	<b>4.8E–08</b>	<b>1.8E–05</b>	2.5E–02	7.6E–01
	KANSL1-AS1	17: 44270942–44274089	27	0.27	0.43	<b>3.4E–10</b>	<b>1.4E–06</b>	2.5E–02	7.6E–01
	LRRC37A	17: 44370099–44415160	75	0.27	0.20	<b>7.9E–08</b>	<b>2.5E–05</b>	9.7E–02	4.5E–01
	LRRC37A2	17: 44588877–44630815	130	0.37	0.18	<b>3.1E–07</b>	8.5E–05	8.4E–02	5.8E–01
q22	STXBP4	17: 53062975–53241646	609	0.20	0.01	<b>1.4E–25</b>	<b>1.6E–25</b>	3.3E–02	<b>1.5E–06</b>

(Continues)

TABLE 1 (Continued)

Cytoband	Gene	Chromosome: Position (start-end)	Number of SNPs	Heritability	Cross validation $r^2$	TWAS $p$ values		
						Overall vs. controls	ER+ vs. controls	ER- vs. controls
q13.2	ZNF404	19: 44376515–44388203	445	0.31	0.06	2.0E–13	5.1E–12	1.7E–08
	ZNF155	19: 44472014–44502477	486	0.39	0.11	8.8E–09	1.0E–07	2.0E–05
	RP11-15A1.7	19: 44501048–44506988	477	0.29	0.14	9.7E–12	7.3E–10	5.1E–07
q11.23	CPNE1	20: 34213953–34220170	299	0.25	0.20	5.3E–05	3.3E–04	1.4E–01

Note: Significant associations after Bonferroni adjustment ( $p < .05/901$ ) are in bold.

Abbreviations: ER, estrogen receptor; SNP, single nucleotide polymorphisms; TWAS, transcriptome-wide association studies.

colocalization, not the effect of modeled gene's expression levels. This additional filter narrowed the number of candidate genes to 901.

## 2.4 | Transcriptome-wide association study (TWAS) analyses

Using the functional weights of those 901 genes and summary level GWAS data, we assessed the association between predicted gene expression and breast cancer risk. We performed summary-based imputation using the ImpG-Summary algorithm (Pasaniuc et al., 2014). Briefly, let  $Z$  be a vector of standardized association statistics ( $z$  scores) of SNPs for a trait at a given *cis* locus,  $\Sigma_{s,s}$  be the LD matrix from reference genotype data and let  $W = (w_1 w_2 w_3 \dots w_j)$  be the weights from the expression prediction model precompiled using the reference panel. Under the null hypothesis that none of the SNPs with  $w_i \neq 0$  is associated with disease, the test statistic  $wZ / (w \Sigma_{s,s} w')^{1/2}$  follows a normal distribution with mean = 0 and variance = 1. To account for finite sample size and instances where  $\Sigma_{s,s}$  was not invertible, we adjusted the diagonal of the matrix using a technique similar to ridge regression with  $\lambda = 0.1$ .

## 2.5 | Case-only TWAS

To assess whether genetically predicted expression was differentially associated with ER+ and ER– breast cancer, we applied the TWAS procedure described above to the  $Z$  statistics from the BCAC case-only analysis. Following arguments in Barfield et al. (2018) the standard TWAS statistic applied to a case-only GWAS results tests hypothesis  $H_0: \beta_2 - \beta_1 = 0$ . This is similar to a conventional multinomial logistic model for subtype-specific breast cancer risk, with expression log odds ratio  $\beta_2$  for ER– disease and  $\beta_1$  for ER+ disease, under which scenario, the expression log odds ratio comparing ER– to ER+ cases is  $\beta_2 - \beta_1$ .

## 2.6 | Conditional analyses

Colocalization makes the interpretation of TWAS hits challenging (Mancuso et al., 2019; Wainberg et al., 2019). In addition to the main TWAS analysis, we also performed conditional and joint (COJO) multiple-SNP analysis at each TWAS significant gene location to distinguish colocalization, and to identify gene(s) independently responsible for the statistical association at each locus. COJO approximates the results of a joint conditional analysis including predicted expression levels from multiple proximal genes. The original COJO approach was designed to assess the association of



individual SNPs with a phenotype; we used an extension that jointly models the associations between multiple linear combinations of individual SNPs (Gusev et al., 2016). We conducted two types of COJO: (a) For regions in which multiple associated features were identified (within 500 kb of each other, i.e., colocalization), we jointly modeled these significant TWAS genes to determine the strongest associated gene (or infer independent signals); (b) To provide information on whether the TWAS gene was responsible for the observed SNP-trait association, we also evaluated whether the GWAS-identified index SNPs remained significant after conditioning on the genes within the same region.

### 3 | RESULTS

#### 3.1 | Breast cancer TWAS

We selected 12,696 transcripts from the 67 GTEx breast tissue samples of European-ancestry women that passed quality control. Based on GCTA-REML analysis, breast-tissue expression levels for 1,355 of these genes were heritable ( $p$  value for  $cis-h_g^2 < .01$ ). We then built linear predictors for these heritable genes and estimated prediction  $r^2$  using fivefold cross-validation. A total of 454 genes failed our cross-validation  $r^2$  requirement ( $r^2 > .01$ ), and we performed TWAS on the remaining 901 genes. We defined statistical significance for TWAS results as a marginal  $p < 5.5 \times 10^{-5}$  (Bonferroni correction controlling the familywise error rate at  $\leq 0.05$  for the 901 genes).

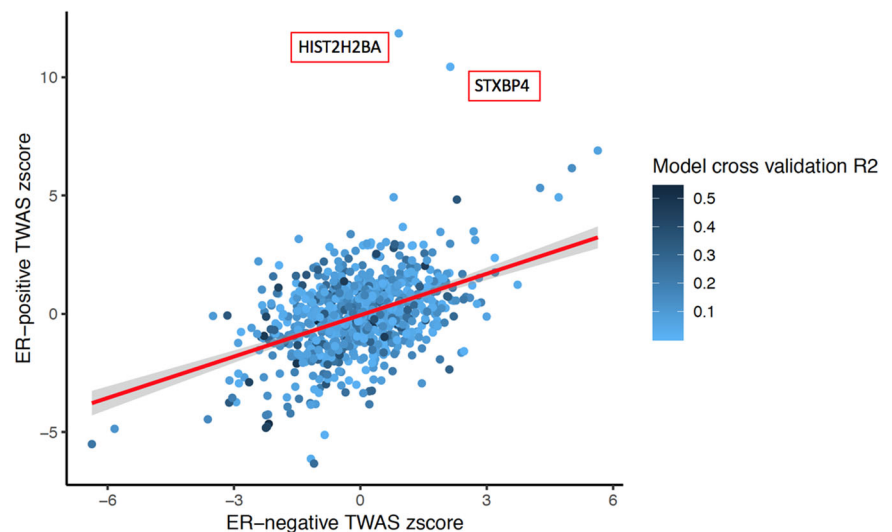
First, to compare with previous GWAS findings and to demonstrate the validity of our results, we performed TWAS analysis in overall breast cancer. We identified 30 genes in 18 cytoband regions associated with breast

cancer risk (Table 1). Of these regions, 11 (containing 21 genes) were previously reported breast cancer susceptibility loci (harboring one or more GWAS-significant SNP). Five genes in the remaining seven regions were previously reported in TWAS or EUGENE analyses (*LINC00886*, *CTD-2323K18.1*, *MAN2C1*, *NUP107*, and *CPNE1*), while the remaining four genes in these regions were novel (*MAEA*, *GDI2*, *ULK3*, and *HSD17B1P1*). *NUP107* and *CPNE1* did not pass a stringent Bonferroni significance threshold in Wu et al. (2018) but passed a less-stringent false discovery rate threshold.

We also carried out analyses focusing on breast cancer subtypes. We found 20 genes associated with ER+ breast cancer, and six genes associated with ER- breast cancer ( $p < .05/901 = 5.5 \times 10^{-5}$ ; Table 1). In our results, all genes associated with ER- disease were also associated with ER+ disease, as well as with overall breast cancer risk. Using a more stringent threshold on the strength of the genetic predictor for expression (cross-validation  $r^2 > .1$ ; 383 genes passed this threshold), we found four TWAS significant ( $p < .05/383 = 1.3 \times 10^{-4}$ ) genes for ER- disease, 14 genes for ER+ disease, and 19 genes for overall breast cancer (18 out of 19 genes are included in Table 1 except for one gene, *CTD/3110H11.1*). As before, these gene sets were nested within each other.

#### 3.2 | Difference of TWAS signal across breast cancer subtypes

We tested whether the imputed gene expression-breast cancer associations differed by subtype using GWAS summary statistics from a case-only analysis, which specifically compared ER+ with ER- breast cancer patients (see Section 2 for details), scanning through



**FIGURE 1** Scatter plot comparing the transcriptome-wide association study z scores in ER+ and ER- patients



901 eligible genes. Two genes, *HIST2H2BA* and *STXBP4*, showed significant associations ( $p < .05/901$ ) with ER status among cases (Figure 1). These two genes were associated with ER+ breast cancer but not associated with ER- breast cancer.

### 3.3 | GWAS signal conditioning on TWAS gene expression

As shown in Table 2, 21 (of 30) TWAS-significant genes were located near GWAS signals. To examine

whether the observed GWAS signal within the gene region could be explained by the expression of that gene, we performed additional analyses conditioning SNP-cancer associations on the predicted expression of that particular significant TWAS gene (See Section 2 and Figure S1, for details). We found that for most regions, GWAS SNPs were no longer associated with the risk of breast cancer once conditioned on the expression of TWAS gene in the region: 15 of 21 genes had no SNPs with a conditional GWAS  $p$  value smaller than the genome-wide significant threshold ( $5 \times 10^{-8}$ ). Thus, there were six genes for which the

**TABLE 2** Summary of conditional analysis at known breast cancer risk region

Gene	Before conditional analysis			After conditional analysis				Magnitude of change in the minimum $p$ value before and after COJO
	Number of SNPs	Number of significant SNPs	Index GWAS SNP $p$ value	Number of significant SNPs	Index SNP	Smallest conditional $p$ values	Ratio <sup>a</sup>	
ALS2CR12	480	12	8.2E-17	1	rs3769823	6.40E-09	0.92	1.28E-08
ATG10	619	24	6.9E-13	0	rs891159	1.20E-07	1.00	5.75E-06
ATP6AP1L	581	24	6.9E-13	0	rs891159	1.70E-07	1.00	4.06E-06
CASP8	493	12	8.2E-17	6	rs3769823	3.90E-12	0.50	2.10E-05
CRHR1	229	13	1.5E-10	0	rs17763086	1.40E-05	1.00	1.07E-05
CRHR1-IT1	230	13	1.5E-10	0	rs17763086	9.90E-05	1.00	1.52E-06
HIST2H2BA	202	19	3.5E-52	1	rs11249433	7.40E-24	0.95	4.73E-29
KANSL1-AS1	34	13	1.5E-10	0	rs17763086	1.60E-01	1.00	9.38E-10
L3MBTL3	724	13	1.7E-12	0	rs6569648	1.40E-03	1.00	1.21E-09
LRRC37A	285	13	1.5E-10	0	rs17763086	4.60E-05	1.00	3.26E-06
LRRC37A4P	285	13	1.5E-10	0	rs17763086	4.60E-05	1.00	3.26E-06
MRPL23-AS1	557	36	2.4E-33	18	rs569550	1.20E-29	0.50	2.00E-04
NUDT17	112	17	1.5E-10	0	rs36107432	3.90E-05	1.00	3.85E-06
RP11-15A1.7	594	32	1E-16	0	rs10426528	4.60E-07	1.00	2.17E-10
RP11-250B2.5	503	8	2.7E-09	0	rs9343989	1.00E-03	1.00	2.70E-06
RP11-554A11.9	532	36	2.8E-44	33	rs680618	2.80E-44	0.08	1.00E+00
RP11-73O6.3	665	13	1.7E-12	0	rs6569648	1.70E-03	1.00	1.00E-09
STXBP4	687	46	2E-28	0	rs244353	2.10E-04	1.00	9.52E-25
ZNF155	597	32	1E-16	5	rs10426528	2.00E-09	0.84	5.00E-08
ZNF404	551	32	1E-16	0	rs10426528	5.30E-07	1.00	1.89E-10
LRRC37A2	152	2	2E-08	0	rs199498	1.80E-02	1.00	1.11E-06

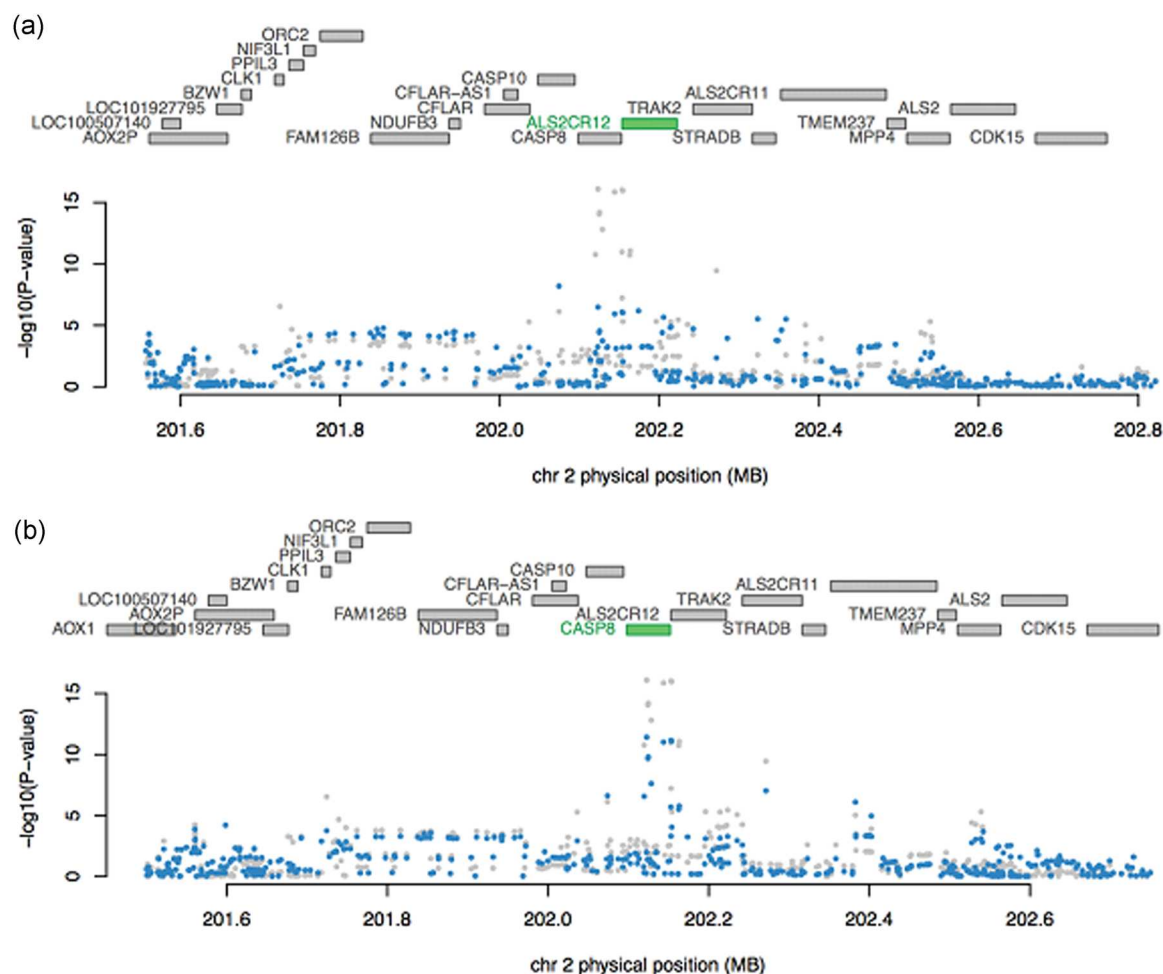
Abbreviations: COJO, conditional and joint; GWAS, genome-wide association studies; SNP, single nucleotide polymorphisms.

<sup>a</sup>Proportion of marginally significant SNPs that are not significant in conditional analyses. Analysis was performed using GWAS summary statistics of ER+ subtypes. The difference between marginal SNP tests for association (GWAS  $p$  values) and the SNP  $p$  values conditional on significant TWAS genes provides some evidence regarding the independence of the TWAS and single-SNP association signals. The number and proportion of SNPs that are genome-wide significant before and after conditioning on a TWAS-significant gene summarizes the degree single-SNP associations are dependent on (or independent of) the TWAS association.

GWAS SNP remained significantly associated with breast cancer risk at the genome-wide threshold ( $5 \times 10^{-8}$ ) after conditioning on TWAS gene expression. The region containing *HIST2H2BA* had only one genome-wide significant SNP remaining, and the region containing *ZNF155* and *ZNF404* had five genome-wide significant SNPs remaining, indicating that the expression of identified genes might explain some but not all of the SNP-breast cancer associations in these regions. For *CASP8* and *MRPL23-AS1* regions, half of the GWAS hits remained genome-wide significant, and for the *RP11-554A11.9* region, 33 out of 36 GWAS SNPs remained (Figures 2 and S1). These results suggest that the genetic association between breast cancer risk and those regions may not be mediated by transcriptional regulation of the genes on which we conditioned.

### 3.4 | Mutually adjusting for TWAS-significant genes in the same region

As shown in Table 3, we identified six regions with more than one TWAS-significant gene: 2q33 (*CASP8*, *ALS2CR12*), 5q14 (*ATG10*, *ATP6AP1L*), 6q22 (*RP11-73O6.3*, *L3MBTL3*), 15q24 (*ULK3*, *MAN2C1*, *CTD-2323K18.1*), 17q21 (*LRRC37A4p*, *CRHR1-IT1*, *CRHR1*, *KANSL1-AS1*, *LRRC37A*, *LRRC37A2*), and 19q13 (*ZNF404*, *ZNF155*, *RP11-15A1.7*). After mutually conditioning on the predicted expression of all significant genes in the same regions, ten genes remained nominally significant ( $p < 0.05$ ). For some regions, only one gene remained, that is *ATG10* for 5q14, *L3MBTL3* for 6q22 and *CRHR1-IT1* for 17q21 (Figures 3a and S2); while for other regions, multiple genes remained significant, including *CASP8* and *ALS2CR12* for 2q33,



**FIGURE 2** Conditional and joint analysis (COJO) for genes near a strong breast cancer GWAS hit. (a) COJO results adjusting for predicted expression of *ALS2CR12*. After conditioning on *ALS2CR12*, almost all original significant GWAS signals (grey dots) disappear (blue dots). (b) COJO results adjusting for the predicted expression of *CASP8*. After conditioning on *CASP8*, some of the original GWAS significant signals (grey dots) remains (blue dots)

**TABLE 3** Conditional and joint analysis of gene region with multiple TWAS significant genes

Region <sup>a</sup>	Gene (colocalized)	Marginal TWAS		COJO	
		Z score	p Value	Z score	p Value
2q33	<b>ALS2CR12</b>	6.7	2.15E-11	4.6	3.70E-06
	<b>CASP8</b>	-5.22	1.76E-07	-2	5.00E-02
5q14	<b>ATG10</b>	-6.37	1.85E-10	-6.37	1.85E-10
	ATP6AP1L	-5.18	2.25E-07	-0.85	0.4
6q22	RP11-73O6.3	-6.92	4.46E-12	0.18	0.86
	<b>L3MBTL3</b>	-7.46	8.45E-14	-7.46	8.45E-14
15q24	<b>ULK3</b>	-4.11	3.87E-05	-4.1	3.90E-05
	<b>MAN2C1</b>	-4.95	7.37E-07	-5	7.40E-07
	CTD-2323K18.1	-4.95	7.49E-07	-1.7	0.083
17q21	LRRC37A4P	6.29	3.12E-10	0.25	0.8
	<b>CRHR1-IT1</b>	-6.3	2.91E-10	-6.3	2.91E-10
	CRHR1	-5.46	4.84E-08	-0.28	0.78
	KANSL1-AS1	-6.28	3.37E-10	-0.04	0.97
	LRRC37A	-5.37	7.89E-08	1.83	0.07
	LRRC37A2	-5.12	3.07E-07	1.81	0.07
19q13	<b>ZNF404</b>	7.35	2.04E-13	3.5	0.001
	<b>ZNF155</b>	5.75	8.81E-09	-2	0.042
	<b>RP11-15A1.7</b>	6.81	9.67E-12	2.8	0.005

Abbreviations: COJO, conditional and joint; TWAS, transcriptome-wide association studies.

<sup>a</sup>Bolded genes remain significant in conditional analyses. Analysis was performed using GWAS summary statistics of ER+ subtypes. Our primary goal in these analyses is to establish whether any of the marginally significant TWAS genes remains significant after conditioning for the most significant gene in the region; since all of the regions with multiple significant genes contain 2–3 significant genes, using a conditional *p* value threshold of .05 is a reasonable threshold for identifying independent signals.

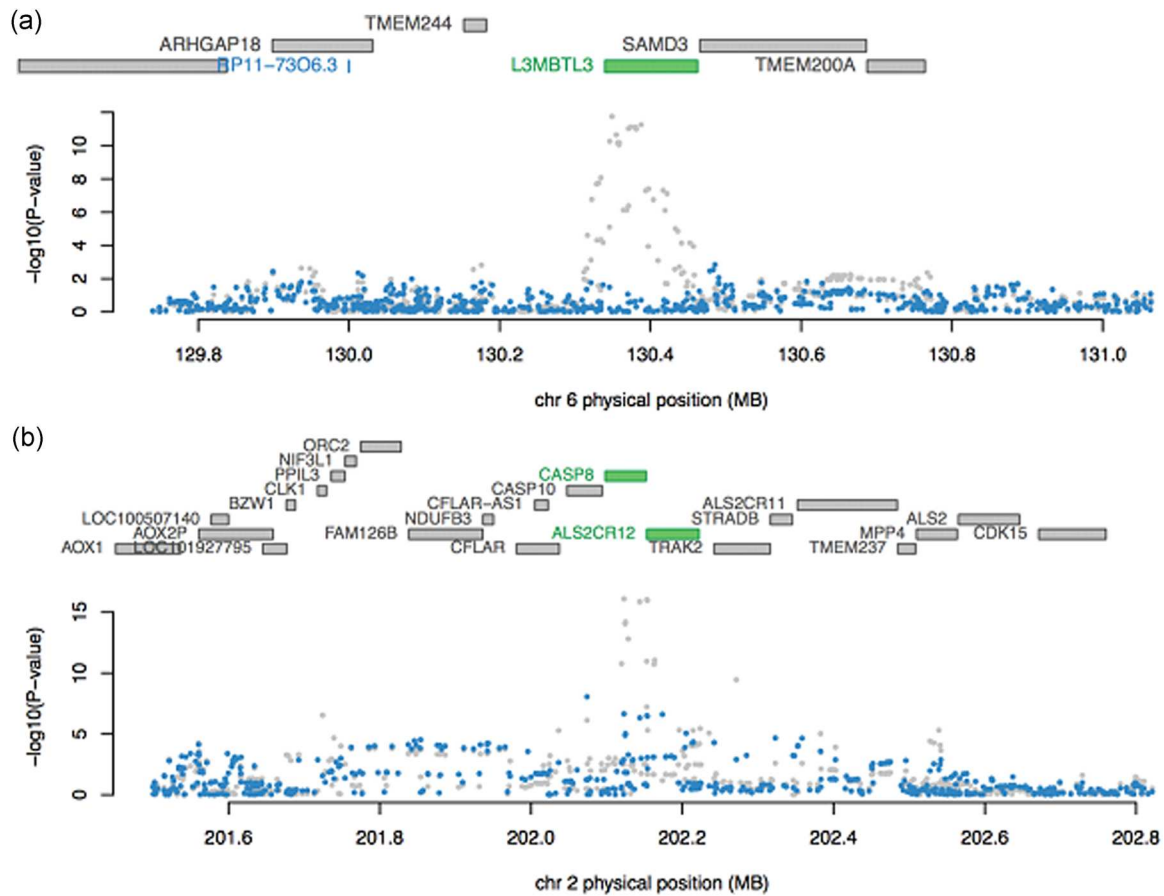
*ULK3* and *MAN2C1* for 15q24, and *ZNF404*, *ZNF155*, and *RP11-15A1.7* for 19q13 (Figures 3b and S2).

## 4 | DISCUSSION

We conducted a TWAS analysis using GTEx mammary tissue gene expression data and GWAS summary data from the largest meta-analysis for breast cancer risk. We assessed associations between overall breast cancer risk and ER+ versus ER- disease. We found 30 genes significantly associated with overall breast cancer risk, 20 genes associated with the ER+ subtype, and six genes with the ER- subtype.

These results are consistent with previous reports from TWAS or similar gene-based approaches, which used various algorithms to build gene expression models. For example, of the 30 genes that we found significantly related to overall breast cancer risk, 23 were also significant in Wu et al. (2018) with very similar test statistics (correlation = 0.96 for the *z* scores between our and Wu's results), and six were significant in Ferreira et al. (2019). One of the six genes we classified as significantly

associated with ER- breast cancer was also found significantly associated with ER- breast cancer in Ferreira et al. (2019). Among these studies, the approach taken by Wu et al. was the most similar to ours. Only seven of the 30 genes that we identified were not identified by Wu et al. (2018), probably due to different *cis*-SNP selection criteria and different candidate genes selected for testing. We defined *cis*-SNPs using a 500 KB window around the gene boundary and included only candidate genes with a significant heritability, while Wu et al. used a 2 MB *cis*-SNP window and included genes with a prediction performance of at least 0.01 without heritability filtering. For genes whose expression could not be predicted well, Wu et al. built models using only SNPs located in promoter or enhancer regions. Despite these methodological differences, the two TWAS results were highly concordant. However, we did not replicate any of the findings in Hoffman et al. (2017) and Gao et al. (2017), which may reflect the smaller sample size of the breast cancer GWAS used in their analyses (3,370 cases and 19,717 controls in Hoffman et al.; 10,597 overall breast cancer cases, 3,879 ER- cases and 11,358 controls in Gao et al.). Specifically, three of the previously reported genes were excluded by our stringent



**FIGURE 3** COJO for regions with multiple TWAS associations. For each plot, the top panel shows all genes in the locus. After COJO analysis, the marginally associated genes are highlighted in blue, while those that remain jointly significant are highlighted in green (in this case, *L3MBTL3*, *CASP8*, and *ALS2CR12*). The bottom panel shows a Manhattan plot of the GWAS signals before (gray) and after (blue) conditioning on the significant (green) genes. (a) COJO results for 6q22 (only one gene remains significant after COJO). (b) COJO results for 2q33 (an example of multiple genes remaining jointly remain significant after COJO). COJO, conditional and joint analysis; GWAS, genome-wide association studies; TWAS, transcriptome-wide association studies

QC procedure (*DHODH*, *ANKLE1* from Hoffman et al. and *TP53INP2* from Gao et al. were not heritable in our analysis) and one was not significant in our analysis (*RCCD1* from Hoffman et al.  $p = .0032$  for overall breast cancer). Both Hoffman et al. and Gao et al. used GWAS results based on a mixed population of European, African, and Asian ancestry (which shared a small set of European samples with our GWAS:  $N < 5,700$  individuals from CGEMS and the BPC3, less than 2% of our GWAS sample). They also used different tissues to build their prediction weights: overall breast tissue (men and women combined, all ethnicities) and whole blood tissue (men and women combined, European ancestry).

Of the 30 genes associated with breast cancer risk in our study, 21 fell into known GWAS regions whereas nine were not close to any known GWAS hit and were, therefore, considered novel. Of these nine genes, five were identified and discussed in Wu et al. (2018) or Ferreira et al. (2019). The four genes uniquely identified

in the present study were *GDI2*, *HSD17B1P1*, *MAEA*, and *ULK3*, several of which have been reported to play a role in breast tumorigenesis or related biological processes. For example, the expression of *GDI2* has been linked with breast cancer through its contribution to enhanced epidermal growth factor receptor endocytosis (EGFR; de Graauw et al., 2014). *HSD17B1P1* is a pseudo-gene related to *HSD17*, which participates in steroid hormone biosynthesis, metabolism, and signaling pathways potentially related to breast cancer risk (Jakubowska et al., 2010). These findings lend support to our results and suggested that further investigation into the roles of the novel genes identified for breast cancer is required.

We performed several conditional analyses not reported in previous TWAS. We examined the local GWAS signals conditioning on the expression of TWAS genes, to provide a measure of how well the expression level of identified TWAS genes explained the local GWAS signals. For many loci, these genes explained a large proportion

of the local GWAS signals and were thus candidates for downstream experimental validation. We also identified candidate genes driving the statistical associations in regions with more than one TWAS gene (usually also regions with known GWAS risk loci) by jointly modeling multiple nominally significant genes. For example, previous studies have suggested that polymorphisms in *CASP8* are associated with breast cancer risk (Cox et al., 2007), whereas a recent paper has shown that the most significant signal in this region is for the imputed intronic SNP rs1830298 in *ALS2CR12* (telomeric to *CASP8*; Lin et al., 2015). Our results provide clarification on whether *CASP8* or *ALS2CR12* expression were more strongly associated with breast cancer risk, since both genes remained significantly associated with breast cancer risk after conditioning on the expression of the other (the conditional  $p$  value for *ALS2CR12* was  $3.70 \times 10^{-6}$ , whereas the conditional  $p$  value for *CASP8* was .05). Eleven of the 12 GWAS hits disappeared after adjusting for the expression of *ALS2CR12*, while half of the GWAS hits remained after adjusting for the expression of *CASP8*. Therefore, we believe that *ALS2CR12* SNPs have a stronger effect and are associated with breast cancer through *ALS2CR12* expression, while *CASP8* remains an additional independent hit, consistent with the latest fine-mapping results (Lin et al., 2015).

Because the genes found to be associated with ER– disease were also associated with ER+ disease, and these, in turn, were associated with overall breast cancer risk, it is difficult to conclude whether the differences in gene sets are due to distinct mechanisms underlying breast cancer subtypes or due to a lack of statistical power because of the smaller disease subtype sample sizes. To address this question, we further incorporated a case-only TWAS comparing ER+ versus ER– breast cancer. We identified two genes, *STXBP4* and *HIST2H2BA*, associated with ER status, which were significantly associated only with ER+ but not ER– breast cancer. Previous studies supported the link between rs6504950 (a SNP in *STXBP4*) and overall breast cancer risk (Antoniou et al., 2010; Warren Andersen et al., 2013). It has also been hypothesized that the risk allele for the two top breast cancer candidate SNPs, rs2787486 and rs244353, affected gene expression of *STXBP4* (Darabi et al., 2016) and CD4 memory cells (Hnisz et al., 2013). One potential explanation for the association between *STXBP4* and breast cancer risk is that it encodes syntaxin binding protein 4, a scaffold protein. In addition, *STXBP4* functions to stabilize and degrade TP63 isoform (a member of the TP53 tumor suppressor protein family), a biologically plausible candidate cancer susceptibility gene. Similarly, SNPs rs2580520 and rs11249433 upstream of *HIST2H2BA* have been identified as breast cancer susceptibility alleles

in a previous GWAS (Bogdanova, Helbig, & Dörk, 2013). Our results suggest that functional and pathway analyses targeting these two genes are likely to shed new light on the differences in tumorigenesis and progression mechanisms between ER+ and ER– patients.

By building gene expression linear predictors in GTEx breast tissue, our analysis offers a tissue-specific model of gene expression. The gene regulatory mechanisms in female breast tissue are arguably the most suitable for studying breast cancer. Moreover, by restricting our reference population to women of European ancestry, rather than mixing genders and ancestries, the resulting gene expression model was a better match to our breast cancer GWAS summary statistics. By using the largest GWAS meta-analysis currently available, we greatly improved the power compared with previous work by Hoffman et al. (2017) and Gao et al. (2017). Finally, by using case-only GWAS summary statistics, we provided insights into genes associated with breast cancer subtype specific risk compared with Wu et al. (2018) and Ferreira et al. (2019).

Similar to previous work by Wu et al. (2018) and Hoffman et al. (2017), our analyses focused on genetic tools trained using expression from breast tissue, chosen because of its direct relevance to breast carcinogenesis. However, given the relatively small sample size in the breast tissue eQTL panel, this choice limited both our power to detect genes with *cis*-heritable expression and the precision of estimated genetic predictors for heritable transcripts. The genetic regulation of expression is constant across tissues for many genes, suggesting that considering other tissues with larger eQTL sample sizes or combining eQTL evidence across tissues may improve power. In addition, other tissues may be relevant for breast cancer development. For example, considering that obesity and hormonal signaling have been linked to breast cancer risk (Bertolini, 2013), gene expression in adipose tissue and brain tissue may have parallel involvement with breast cancer etiology. We are currently developing methods for cross-tissue TWAS, using sCCA (sparse canonical correlation analysis) to build features that combine gene expression values across tissues that share similar genetic regulation mechanisms, while allowing tissues with different regulation patterns to contribute to different features (Feng, Pasaniuc, Major, & Kraft, 2018).

In conclusion, we have identified new breast cancer target genes both for functional experiments and as causal gene candidates in the significant TWAS gene regions. We have also identified associations between gene expression and breast cancer risk specific to disease subtypes, where two novel genes have been found specifically associated with ER+ breast cancer risk.



This analytic strategy warrants application in studies aimed at defining the genomic architecture of cancers other than breast cancer.

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## CONFLICT OF INTERESTS

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## DATA AVAILABILITY STATEMENT

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## SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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